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**Endoluminal high-resolution MR imaging protocol for colon walls analysis in mouse
model of colitis**

Hugo Dorez¹, Raphaël Sablong¹, Laurence Canaple², Hervé Saint-Jalmes^{3,4}, Sophie Gaillard¹,
Driffa Moussata^{1,5} and Olivier Beuf¹

¹Université de Lyon, CREATIS; CNRS UMR 5220; INSERM U1044; INSA-Lyon,
Université Lyon 1, Villeurbanne, France

²Institut de Génomique Fonctionnelle de Lyon, Université de Lyon 1, UMR 5242 CNRS,
Ecole Normale Supérieure de Lyon, Lyon, France

³LTSI; INSERM U642; Université Rennes 1, Rennes, France

⁴CRLCC; Centre Eugène Marquis, Rennes, France

⁵Hôpital Régional Universitaire de Tours - Service hépato-gastroentérologie, Tours, France

SHORT TITLE: Endoluminal MR imaging protocol for mouse colon walls exam

Corresponding author:

Olivier Beuf, PhD

CREATIS, CNRS UMR 5220, Inserm U1044

Université Lyon 1, bât. 308 (ESCPE)

3 rue Victor Grignard

69616 Villeurbanne, FRANCE.

Tel: +33 4 72 43 15 97

Fax: +33 4 72 44 81 99

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40 ABSTRACT:

41
42 Object: An endoluminal MR imaging protocol including the design of an endoluminal coil
43 (EC) was defined for high spatial resolution MR imaging of mice gastrointestinal walls at
44 4.7T.

45 Materials and Methods: A receive-only radiofrequency single-loop coil was developed for
46 mice colon wall imaging. Combined with a specific protocol, the prototype was first
47 characterized *in vitro* on phantoms and on vegetables. Signal-to-noise ratio (SNR) profiles
48 were compared with a quadrature volume birdcage coil (QVBC). Endoluminal MR imaging
49 protocol combined with the EC was assessed *in vivo* on mice.

50 Results: SNR measured close to the coil is significantly higher (10 times and up to 3 mm of
51 the EC center) than the SNR measured with the QVBC. The gain in SNR can be used to
52 reduce the in-plane pixel size up to $39 \times 39 \mu\text{m}^2$ (234 μm slice thickness) without time penalty.
53 The different colon wall layers can only be distinguished on images acquired with the EC.

54 Conclusion: Dedicated EC provides suitable images for the assessment of mice colon wall
55 layers. This proof of concept provides gain in spatial resolution and leads to adequate
56 protocols, for the assessment of human colorectal cancer that can now be used as a new
57 imaging tool for a better understanding of the pathology.

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59
60 KEYWORDS: Colorectal Neoplasms, Magnetic Resonance Imaging, Colitis, Instrumentation

INTRODUCTION:

Every year, on the overall cases of cancer diagnosed in the world, 13% are colorectal cancer (CRC) [1–3]. It is the second deadliest cancer in Western countries just behind lung cancer. The 5-years survival rate is higher than 80% when the cancer is diagnosed and treated at an early stage [4]. The cancerogenesis of CRC has not been yet precisely established but two major pathways have been identified [5]: the chromosomal instability pathway (adenoma-carcinoma sequence) and the microsatellite instability. Those pathways induce changes in the cellular proliferation with abnormal cells growing and infiltration into the gastrointestinal walls. The different stages of CRC infiltration are well known [6, 7]. It is first characterized by an inflammation stage affecting the superficial layer of the colon wall. Then, the abnormal cells proliferation infiltrates the wall to deeper structures and, eventually, reaches the lymphatic system. Because the survival of patients strongly depends on the diagnosis stage, an early and accurate diagnosis to improve the therapeutic response efficacy is highly beneficial to the patient. Magnetic Resonance Imaging (MRI), computed tomography (CT), positron emission tomography (PET) and some optical modalities provide different kind of information when imaging gastrointestinal walls. However, it is still difficult (using those modalities and associated protocols) to assess fine structures from the abnormal cells proliferation. Today, endoscopic ultrasound (EUS) is widely used in clinical routine for the staging of colorectal cancer lesions [8–10]. It is an available and low cost real-time (motion free) imaging modality. Moreover, the sensitivity of EUS for rectal tumors invasion is approximately 88-95% while specificity is nearly 100% (for stage 2 and more) [11]. However, EUS is especially used on rectal carcinomas and fewer studies have shown the potential of this modality on colon cancer [12–14]. EUS is also strongly dependent on operator and patient and thus require practice to perform an accurate diagnosis. This is especially the case when examination is

performed at early stages of CRC. Some studies describe the use of endoscopic MRI compared to EUS [15–19] to assess colorectal region. MRI is well suited to investigate human soft tissues and provides a good spatial resolution for the characterization and staging of gastrointestinal lesions [20]. The modality provides ways of performing high-resolution quantitative maps, various contrast imaging and three-dimensional acquisitions but also performing magnetic resonance spectroscopy. However, colon and rectal walls assessment requires an even higher spatial resolution for visualizing different wall layers in detail [21]. It has been shown that spatial resolution in MRI could be used to depict more precisely, using EC, the gut wall than EUS, for N-staging [17] for example or for the staging of inflammation degrees and cancer lesions. Despite the fact that endoscopic MRI has demonstrated to be superior to MRI using external array coils, by providing exquisite information of wall structures [16, 19, 22, 23], the use of endoscopic coils in human can induce strong local temperature increase at coil tip and along the coaxial cable [24–26]. While heating can be strongly reduced adding for example radiofrequency traps [24, 27], the use of inner coils on humans was currently limited to prostate or rectum and endoscopic MRI of colon based on the use of endoluminal coil is not performed on patient. Especially due to the Human colon anatomy (angle at the rectosigmoid junction) and the invasive procedure (requires anesthesia and insertion of the coil into the patient's rectum).

Today, a better understanding of the CRC genesis to adapt therapeutics responses and increase the survival of patients is needed. This goes through the following of the adenoma-carcinoma sequence (inflammation to cancer) on a model of CRC. The follow-up of this sequence on Human longitudinal studies based on endoscopic MRI is not yet possible due to current safety issues. Moreover, the development of CRC on Human is a long process that can last over decades to understand the growth mechanisms (inflammation evolving into cancer). Nevertheless, this can be done on a mouse model of colitis [28], chemically induced by the

combination of dextran sulfate sodium (DSS 2% in drinking water) and azoxymethan (AOM, 10mg/kg). Trying to understand the growth mechanisms of CRC and to assess the potential of MRI in a mouse model of colitis present several advantages: first, the pathology induced by the method introduced by *Tanaka et al* in 2003 is a reliable model with strong correlations between Human and mice CRC; then, it is possible to follow mice on a much shorter time period (typically six months); and, finally, the examination can be performed for each stages. This paper is describing an endoluminal MRI protocol dedicated to the examination of a mouse cancer model as a preliminary and mandatory step before to move to clinical Human CRC examinations.

While MRI is well suited to investigate abdominal-anatomical structures [20], colon and rectal walls assessment requires an even higher spatial resolution for visualizing different wall layers in detail [21], even more on mice (colon wall thickness 0.3-0.7 mm [29]), still within a limited acquisition time suitable for *in vivo* examination.

Endoluminal Coil (EC) is drastically increasing the signal-to-noise ratio (SNR) close to the coil compared to external array coil [16, 19, 22, 23]. This kind of elongated loop, based on two coplanar copper traces, was already used for intravascular MRI and MRS in the mid-nineties [30]. The temperature around the EC was measured during *in vitro* acquisition [31]. No abnormal and critical temperature increasing was observed. EC can provide a higher spatial and/or temporal resolution and image quality to improve the characterization and staging of colorectal lesions. The aim of this study was to develop, build and characterize an EC for mice rectal wall investigation together with the adapted protocol. The proof of concept of the whole protocol was demonstrated on a limited number of examinations at different stages of a mouse model of CRC.

MATERIALS AND METHODS:

Endoluminal Coil description:

The EC was designed for mice colon investigation on a 4.7 T Bruker Biospec system (Bruker, Ettlingen, Germany). It is a single loop with dimensions sized to the colon mouse anatomy [32] which leads to a 30 mm length and a 1.6 mm outer diameter (heat shrink coating included). The copper tracks are 0.8 mm width and 0.8 mm thick with a copper thickness of 35 μm . Following these characteristics, the inductance and resistance of the copper tracks were calculated using the analytical expression of two-distant traces equation. The coil loop and tuning and matching circuit (including a ground plane on the back side of the circuit) were mechanically carved (Circuit Board Plotters S63, LPKF Laser and Electronics®, Germany) on both sides of a FR4 epoxy substrate of 0.8 mm thickness (C.I.F®, Buc, France). Nonmagnetic electronic circuit components were soldered on a single face. The two longest conductors of the loop have been placed on each side of the substrate. A copper tape was used at the distal end of the coil to close the loop between the two faces (Fig1). One for connecting the loop to the upper face (straight arrows) and, the second one, to connect the direct current (DC) ground and radiofrequency (RF) signal ground together (dashed arrows) to the ground plane. This is done to avoid current loops between two different grounds. The RF signal was carried out by a coaxial cable (RG-58, Radiall®, France) and the DC current travels through a 8-wires cable (Alpha Wire®, New Jersey, USA) connected on a non-magnetic Fisher connector (Plugs 4032, Fisher Connectors®, Suisse) to the coil. This latter decoupling cable is connected to the 14 pins connector of the Bruker decoupling box cable through a homemade Fisher-Bruker conversion cable. A heat shrink coating (PF3-135) houses the loop coil to isolate the tracks from colon and rectum tissues. The end of the coil was sealed with glue (Araldite PRECISION, Araldite®, Bâle, Switzerland). When the EC is used, the

electrical circuit is isolated with polytetrafluoroethylene (PTFE or Teflon, DuPont®, Wilmington, USA) to avoid contacts with tissues or liquids.

Endoluminal Coil characterization:

The resonant circuit is based on a RLC resonator circuit made of fixed nonmagnetic components (ATC, Huntington Station, New York, USA) and trimmer components (AT 57290, Temex Ceramics®, Bordeaux, France) – Fig 1. Considering only the intrinsic parameters of the coil, which are the coil geometry, the materials used and the coupling effects (mutual inductance for example); the loop inductance and resistance were estimated to 33.2 nH and 0.1 Ohm respectively. The values were calculated and corrected using measurements on a Vector Network Analyzer (E5070B – ENA Series, Agilent Technologies®, California, USA) - VNA. Intrinsic characteristics of electronic components (series resistances, self-inductances, parallel capacitances, self-frequency resonance) were modeled to simulate the precise electronic behavior of the coil. Modeling was done using a Simulation Program with Integrated Circuit Emphasis (LTspiceIV, Linear Technology®, Milpitas, USA) feed with electronic components characteristics based on their individual datasheets. The value and manufacturers of the electronic components are summarized in table 1.

In the coil design, there are basically four functions to achieve. Briefly, the capacitor C_t insures the tuning function, changing this value adjust the resonance frequency of the coil (on the S11 magnitude chart). Capacitors $C_{m,1}$ and $C_{m,2}$ are used for matching the coil to the 50 Ohms impedance of receiver circuit in order to maximize the signal power transmission. The $C_{B,1}$ and $C_{B,2}$ capacitors are used for blocking the DC current into the loop coil to improve the decoupling efficiency. It could also create a local static magnetic field interfering with the static magnetic field B_0 . This ensures the correct decoupling of the loop coil. Mounted in

series with C_t , capacitors can also contribute marginally for tuning the coil. Trimmer capacitors can be tuned mechanically with a nonmagnetic screwdriver to adjust the coil to match to 50 Ohms at the working frequency of 200 MHz (proton resonance frequency at 4.7 T). Finally, the decoupling circuit is represented by two Choke inductors and one PIN diode. The Choke inductors $L_{B,1}$ and $L_{B,2}$ protect the receiver amplifier from the induced RF current during the RF transmit pulses.

The PIN diode was mounted in parallel to the loop coil to minimize the series resistance and so maximizes the Quality factor (Q-factor) and thus the SNR [33, 34]. The quality factor of the coil was measured using a Vector Network Analyzer (VNA). The loaded Q-factor Q_L was measured with the EC inserted into a phantom containing a solution of saline solution and the unloaded Q-factor Q_0 was measured without the phantom.

SNR assessment:

All the experiments were performed on a 4.7 T Bruker Biospec System (Bruker, Ettlingen, Germany). In order to characterize the EC, each acquisition done with the EC was also compared to a ^1H 32 mm inner diameter quadrature volume birdcage coil (Rapid Biomedical®, Rimpar, Germany) – QVBC dedicated to mouse body. However, the QVBC used does not include any decoupling circuit. As a consequence, the EC should not be inserted within the sample during QVBC operation. That is the reason why, when using the QVBC, an optical fiber (HCG 600, OFS®, Atlanta, USA) surrounded by a heat shrink coating was inserted into the mouse rectum instead of the EC. The outer diameter of the optical fiber and the heat shrink coating is 1.6 mm. Then, when using the EC, the QVBC was replaced by a ^1H 72 mm inner diameter transmit linear volume coil (Rapid Biomedical®, Rimpar, Germany) – VC including an active decoupling circuit. A protocol was developed and optimized for the purpose of EC assessment on phantom and small animals. Three axial 3D fast low-angle-shot

(FLASH) sequences were used for anatomical and structure imaging but also for computing parametric T1 maps [37]. Then, one coronal FLASH 3D Slab sequence is performed to assess the colon along the EC. Finally, for computing parametric T2 maps, a 2D multiple spin-echo sequence was carried out. Table 2 summarizes the sequences parameters used in the *in vivo* protocol.

For SNR gain assessment, the coil was inserted into a 12 mm outer diameter tube filled with a solution of 1.25 g/L of $NiSO_4$ and 5 g/L of $NaCl$ to mimic approximately the loading conditions found in living tissues and have convenient relaxation times (T1-value of 218 ms and T2-value of 140 ms). First, the receiver EC was placed at the center of the VC used in transmit mode. The EC was driven by the active decoupling circuit during transmission. For SNR comparisons, axial images were acquired using FLASH 3D Slab sequences (table 3). Then, VC and EC were replaced by the QVBC and the optical fiber, respectively. With the same protocol, MR images were acquired using the QVBC (emission/reception mode). Slab was positioned manually to scan the same region in order to allow co-localization. The SNR were compared after correcting for difference in voxel size including the antialiasing steps, receive bandwidth and number of excitations between the EC and VCs [38]. SNR profiles were normalized using this method and plotted in the same graph.

Spatial resolution and contrast enhancement using an EC:

The improvement of SNR provided by the EC can be used to decrease the voxel size without acquisition time penalties. To assess the spatial resolution achievable in the region of interest (ROI) with the EC within a reasonable acquisition time, a phantom containing the same $NiSO_4$ solution was filled with optical fibers of standard diameters (ranging from 150 μm to 1000 μm). Using the transmit VC and the EC in reception mode, two sequences were acquired. First, an axial FLASH 3D Slab with a pixel size of 83x83 μm^2 (slice of 234 μm

thick) was carried out(sequence 1 in table 3). Then, the sequence was adjusted to reduce the voxel size up to $39 \times 39 \mu\text{m}^2$ keeping a slice thickness of $234 \mu\text{m}$ (sequence 2 in table 3). The structures observed were measured using the software ImageJ [39] and compared to the optical fiber diameters measured with a digital caliper.

After having characterized the ECs on inorganic phantoms, experiments were carried out on organic phantoms such as onions in order to evaluate the sensitivity of such coils. Onions offer fine lamellar structures divided in several bulbs inside. The thickness of the different layers is, in average, the same as the one expected in mouse colon wall. Images performed on vegetables provide good markers to evaluate the sensitivity of the coil by analyzing the finest structure observable (for a given SNR). Experiments were performed using 3D Slab FLASH sequences (20 mm FOV, 25 mm slab thickness, $\text{TR/TE} = 25/4.31 \text{ ms}$ with 25° flip angle, $256 \times 192 \times 96$ matrix, 36 764 Hz bandwidth and 11'31'' acquisition time). The VC was turned in emission mode and the EC switched on reception mode. Finally, the spatial resolution was increased by reducing the pixel size down to $39 \times 39 \mu\text{m}^2$ (10 mm FOV, 15 mm slab thickness, $\text{TR/TE} = 20/6.1 \text{ ms}$ with 20° flip angle, $256 \times 256 \times 64$ matrix, 25 kHz bandwidth and 10'55'' acquisition time) with a slice thickness equals to $234 \mu\text{m}$.

In vivomouse colon examination:

In vivo experiments were performed to evaluate the feasibility and interest of using an EC in a mouse model of colitis. The study was carried out on mice chemically treated during two months to induce colitis using a combination of azoxymethan (AOM, intraperitoneal injection, 10 mg/kg body weight) and dextran sulfate sodium (DSS, in drinking water, concentration of 2%) and four mice without treatment used as control. The experiments were in accordance with the rules and regulations of the UCBL Ethics Committee on animal experimentation. Prior to the examination, mice were kept on a 12 hours day/night rhythm in

261 a 300 cm² plastic cages with straw bedding, pellet food and tap water. Mice were anesthetized
262 using an isoflurane tabletop station (TEM Sega®, Lormont, France). Animals respiratory
263 index was monitored during the experimentation by using a pressure sensor placed on the
264 mouse chest. During the induction phase, mice were anesthetized with 3% of isoflurane and
265 aspiration flow set up on 0.4 L/min. During imaging, the anesthesia was maintained with
266 1.4 to 1.7% isoflurane vaporization and aspiration flow set up on 0.4 L/min. Images of colon
267 wall were first acquired by using the transmit/receive QVBC. In that case, to mimic the
268 presence of the EC, an optical fiber was carefully introduced into the mouse rectum using
269 sterilized lubricant KY gel (K-Y®, Johnson & Johnson, USA) and held in position with
270 adhesive tape (strap on the MRI bed). Then, to assess colon wall layers with the EC, the
271 QVBC was replaced by the linear VC. MR images were acquired to assess colon wall layers
272 using described protocol in table 2.

273 For each animals, T1- and T2-maps were computed using a linear fit from sequences acquired
274 at different flip angle (15°, 20°, 25°) [37] and an exponential fit [40] at six different echo
275 times (12.8 ms, 25.21 ms, 37.81 ms, 50.42 ms, 63.03 ms and 75.63 ms), respectively. Before the
276 computation, images were threshold with an automatic method. The threshold level was set to
277 three times the standard deviation measured in a ROI with no signal (located in the top left
278 corner of the image far from the EC). Two additional maps, representing the T1 linear fit
279 quality and the T2 exponential fit quality, were also computed. To ensure reliable values of
280 T1 and T2 relaxation times, fits having a coefficient of determination (R^2) inferior to 0.98 were
281 systematically excluded from the representation.

282 For each *in vitro* and *in vivo* experiment, the coil was held in position with adhesive tape and
283 not tilted during acquisitions. The EC was oriented in such a way to keep the B_1 of the EC
284 perpendicular to both static B_0 and transmit RF B_1 magnetic fields. Hence maximum
285 sensitivity and geometric decoupling are ensured respectively. As the colon natural behavior

286 is to fold, the EC is always encompassed by colon structures ensuring almost uniform radial
287 signal intensity.

288

RESULTS:

An endoluminal MR coil, fitting the constraints for *in vivo* experimentation on mice (colon and rectum walls assessment) was designed and built. The prototypes are waterproof, reusable, biocompatible (with the investigated environment) and can be tuned/matched to the appropriate resonance frequency. The measured loaded Q-factor (Q_L) of the coil was 32 and the coupling coefficient was found equal to 1.25. This leads to a value of 72 for the unloaded Q-factor. The adaptation, at the 200.13 MHz working frequency in loading conditions was 50.03 Ohms. The decoupling efficiency of the EC was tested using a VNA. When the pin diode is activated, the resonance frequency is shifted to 150 MHz and the isolation at the resonance frequency reach -43 dB.

The EC provide a higher SNR in the close vicinity of the coil than the one obtained with quadrature volume birdcage coil. The EC geometry and dimensions fit with the use of the volume birdcage coil (72 mm inner diameter), without causing interferences and/or impacting the image quality. The absence of signal highlights in the close vicinity of the EC was confirmed when acquiring images with the body coil. Indeed, the continuously decoupled EC, not impacting the image around loop coil, was attested the correct functioning and efficacy of the decoupling circuit. The results of the SNR obtained with the EC compared with the QVBC acquired on phantom solutions are shown in figure 2. In receive mode, the SNR profile of the EC was much higher than the one achieved with the QVBC up to 3 mm apart from the coil center. SNR profiles plotted in figure 2 were corrected regarding the voxel size, bandwidth and number of excitations for a straightforward comparison. The EC SNR gain is 10 times greater at the close proximity and reaches the SNR of the QVBC at approximately 3 mm from the center.

A precise measurement of the structures observable in the loop-coil field of view was achieved on the phantom constituted with calibrated optical fibers (figure 3). Optical

fibers diameters were measured with a standard deviation of $\pm 6.3 \mu\text{m}$ using the ImageJ software and then compared with a digital caliper (nominal precision $\pm 10 \mu\text{m}$).

Experiments led on vegetables have shown the feasibility to obtain high spatial resolution images and depict structural details. It is possible to distinguish the lamellar structures of the onion to approximately 3 mm of the coil center (figure 4). White arrows on figure 4 localize the same thin slice of the onion obtained with the VC (figure 4a) and with the EC (figure 4b and 4c) at two different spatial resolutions (different SNR).

***In vivo* Experiments**

The EC geometry fits the mouse anatomy and is adapted to *in vivo* experiments. Once the mouse is anesthetized, the insertion of the EC performed by an experimented gastroenterologist is simple and harmless for the animal. The animal's recovery time is short and no signs of prostration and/or pain were observed during the following few hours in accordance with the French legislation. Images acquired *in vivo* show a very good quality (figure 5), in terms of contrast and high spatial resolution. The use of the QVBC does not allow a correct visualization of colon walls due to limited achievable spatial resolution. Whereas the SNR gain provided by the EC is used to increase spatial resolution and to depict nicely the mouse rectum wall layers and surrounding structures. On figure 5b, the map ratio of $\text{SNR}_{\text{EC}}/\text{SNR}_{\text{QVBC}}$ is overlaid on MR images and undoubtedly illustrates the improved SNR area given by the EC. The white arrow on figure 5a1 and 5c locates a blood vessel used for co-localization. The examination with the EC gives a better visualization of colorectal wall layers and surrounding structures (figure 5d and 5e). White arrows locate also the mucosa – submucosa complex. Using such a coil allows to increase spatial resolution up to $39 \mu\text{m}$ in-plane resolution (with $234 \mu\text{m}$ slice thickness, see figure 6a and 6b). It is then possible to well

338 distinguish the mucosa complex in details. ① is the mucosa, ② the submucosa and ③ the
339 muscularis externa. On figure 6b, ④ locates a thin muscle layer.

340 The T1- and T2-values of structures around the EC can be seen on figure 7. The maps have
341 been calculated and plotted with an estimation of the fit quality to ensure reliable values. It is
342 worth noting that this parametric procedure removes the strong signal intensity decay that
343 could lead to misinterpretation on the magnitude images. T1- and T2-maps plotted from two
344 different slices illustrate different structures; ① identifies a muscle structure and ② the colon
345 wall. The averaged T1 relaxation times were measured at $800\text{ ms} \pm 100\text{ ms}$ and
346 $1200\text{ ms} \pm 200\text{ ms}$ respectively. Then, looking at the T2 maps, mucosa layer and muscularis
347 externa are clearly differentiable (③ and ④ on figure 7). The T2 relaxation times are nearly
348 identical, $80\text{ ms} \pm 10\text{ ms}$ for both structures. Finally, * locates the submucosa which tends to
349 have lower T2 values ($50\text{ ms} \pm 10\text{ ms}$). The fit quality maps show a reproducible method for
350 T1- and T2-values with only a few points excluded in the field-of-view considered.

351

DISCUSSION:

EC have been used since the late eighties [41] but essentially for visualizing human prostate and rectum regions [42–45]. Currently, the major problem lies in the heat induced by the RF current into conductors. Several studies have shown the potential of using RF traps to reduce this increase in temperature [24, 26, 46]. Optical transmission and optical decoupling system have also been proposed in order to replace conventional coaxial cable [47, 48]. More recently, the exploration of metamaterials have opened new perspectives for alternative ways of RF signal transmission [49]. These metamaterials could overcome several coaxial cable disadvantages such as high propagation loss or heating along the conductor in the presence of electrical fields. But also, reaching the Human colon is difficult due to the 90° angle and assessing higher structures in the colon may require flexible endorectal MR probes. Other studies have shown the high potential of using EC on preclinical models such as pigs [50], rabbits [16, 19] and rats [15]. But, it is the first reported study assessing colon and rectum walls using dedicated endorectal coils on a mouse model of colitis (inflammation evolving into cancer).

In this study, the feasibility to design an EC for small animal colon wall imaging such as mice was demonstrated. It has been shown that EC can provide typical 40 µm in-plane pixel images with a slice thickness of 234 µm along digestive tract. External volume coils provide less detailed images in similar conditions (same sequence parameters and position). It enables the visualization of colon wall layers and deeper structures with a spatial resolution and SNR not achievable with quadrature volume birdcage coil dedicated to mice gastrointestinal imaging.

For decoupling the EC during the emission of radiofrequencies RF pulses, a parallel decoupling circuit was used. This choice was made to decrease the total resistance of the coil. Indeed, regarding the coil's dimensions and components, one third to one half of the total

377 circuit resistance is due to the PIN diode characteristics alone.Placing the PIN diode in series
378 leads in higher decoupling efficiency but decreases the quality factor and the SNR as well.
379 When the PIN diode is placed in parallel of the tuning capacitor, the resistance is lowered.
380 The quality factor of the coil is improvedbut the decoupling efficiency is reduced. The PIN
381 diode is not entirely shutting down the loop coil and a weak fraction of the radiofrequency
382 signal can still go through into the loop.In principle a coil having one or two parallel PIN
383 diodes mounted in series with the loop would benefit from a better decoupling performance.
384 Even if this solution is feasible for *in vitro* experiments it is not compatible with *in vivo*
385 experiments.PIN diode used here are bigger components than case A ATC capacitors. Several
386 PIN diodes on the coil will result in blocking the access to trimmer capacitors.
387 Using those trimmer capacitors help for tuning/matching the coil at the desired frequency but
388 this kind of component has a poor life time, due to their mechanism working with a
389 screwdriver, and had to be changed regularly. Usually, trimmer capacitors are linked to
390 insulating sticks for tuning and matching. In the case of small coils it is difficult to have such
391 a configuration.

392 A heat shrink coating covers the EC during*in vivo* or *in vitro* experiments to prevent the
393 sensor from immediate short-circuit risk. Consequently,the copper tracks are isolated from the
394 surrounding mediumbut this protection layer weakens the maximum SNR achievable in the
395 close vicinity of the coil. Using a thinner protection could improve the SNR close to the
396 coil.Besides, the EC is wrapped with PTFE to protect the electronic circuit during *in vivo*
397 experiments:as the coil is used in wet environment, long-term oxidation of the copper tracks
398 can appear if the PTFE is not removed after each use and the EC not properly dried.
399 Oxidation could lead to short-circuits, deterioration of the copper tracks and/or components
400 and degradation of transmission line properties.A protective plastic cover will be designed
401 and printed to improve protection. SNR comparisons could not be achieved using the QVBC

at the exact same position due to the absence of decoupling circuit on this coil. The comparison was then done sequentially slice by slice and correlated with images acquired with the EC. Mouse's body examination would hardly be done with the available 70 mm inner diameter birdcage coil. However, the use of a 40 mm inner diameter linear birdcage coil with active decoupling system would easily allow switching between EC and volume coil. Beside the easier and straightforward comparison in SNR, this setup would allow to acquire very local images of the colon as well as more regional images of the mouse. This setup would be a considerable way of improvement.

The phantom containing optical fibers was useful to assess the size of structures that could be observed (from 150 μm to 1 mm) in the EC FOV. On the figure 3, optical fibers seem to be equally distributed around the EC. As optical fibers fell down to the bottom of the container the upper part of the phantom was not containing enough fibers. More of them will have to be added to distribute correctly optical fibers around the sensor. Despite the fact that the sequences chosen for imaging the phantom had a slice thickness of 234 μm , no partial volume effect was observed when measuring outer diameter of the optical fibers.

For the experimentations done on vegetables, onions have been chosen for their structure. Despite the fact that onions present very thin lamellar patterns, it can be assimilated to a semi-rigid material. The molecules of water are constrained by their environment. So, when looking closely to the lamellar structure of this vegetable, it appears that air gaps are located between each slice. This causes some susceptibility artefacts. Moreover, the slice position chosen for the comparison was not exactly the same than the one obtained with the VC. To be able to localize the same structures between body coil images and EC images, the VC was used instead of the QVBC. The SNR of the VC allows the overall visualization of several layers of the vegetable. But to depict lamellar structure, the VC is inappropriate due to insufficient SNR. The use of the EC increases drastically the local SNR and fine structures can be

427 visualized. But it also causes a highlight on the image, in the close vicinity of the coil. To
428 obtain the same details with the QVBC, the same sequence was done using six averages; the
429 acquisition time was increase to 1h15min.

430 *In vivo* acquisitions show a huge improvement in the differentiation of colon wall
431 layerscomparedto the QVBC. The designed EC fits with the straight configuration of the
432 mouse colon. Indeed, human colon presents curves which cannot be passed using such a rigid
433 coil. For one of the healthy mouse, the colon was sampled for further analyzes.During the
434 dissection, the length, average diameter and thickness of the colon were approximately
435 measured, using a digital caliper, to optimize the coil loop.

436 The protocol duration developed for the need of the study wasshort enough (~38 min) to
437 ensure the safe examination and recovery of the animal. It provides anatomical and parametric
438 images (T1 and T2 values). Three scout sequences with orthogonal plane orientationswere
439 first done with the VC in emission/reception mode. Acquisitions were compared between
440 images acquired with and without a trigger deviceusing a respiratory sensor place on the
441 lower part of the abdomen. Though respiratory movement is ample and cardiac beats are
442 rapid, those movements are far from the colon and no significant differences were observed.
443 So, no sequence triggering was performed in order to reduce the acquisition time.
444 Nevertheless, colon can be affected by peristaltic movements.These movements have been
445 evaluated using an optical system developed for this purpose. Movements of colon walls are
446 slow, not regular and limited in amplitude.The degradation induced by this motion on image
447 quality is thus difficult to evaluate.A further analysis will be done to quantify this movement
448 and see the influence on MR images.

449 The EC provides a high SNR in its close vicinity but the SNR drops down heavily. Up to
450 3 mmfrom the center, the SNR is higher than the one with the QVBC, further the sensitivity
451 of the EC begins very poor. The colon wall thickness was measuresin the range of

0.2to0.5 mm (0.3 – 1mm in the literature [28, 51]). Regions of interest, such as colon walls, are closely located to the loop coil where the EC provides a very high SNR. Looking further the two first millimeters is not required for the purpose of this study. On figure 6, the in-plane pixel size was reduced ($\sim 39 \mu\text{m}$ pixel width with a slice thickness of $234 \mu\text{m}$). As the SNR is decreasing with the improvement of spatial resolution, the gain in SNR provided by EC is used to define the best compromise between pixel size and signal intensity. As the QVBC does not possess a decoupling circuit, the colocalization was done using anatomical markers. Blood vessels are convenient to localize the slice position. The mouse bladder and vertebrae can also be used as convenient landmarks.

Even if an optical endoscopic control exam is realized before inserting the coil into the mouse rectum, sometimes the lumen is still filled with feces and other natural substances. Those feces tend to drastically decrease the signal leading in an important loss of contrast around the feces. The systematic endoscopic control combined with a medication controlling the peristaltic movement (loperamide for example) could help to reduce the presence of feces in the lumen. During the evolution of the pathology, bleeding can also appear. When the EC is pulled out from the colon the presence of blood is sometimes noticed. This also induces distortion into the image quality and has to be taken into account for image processing.

Computed parametric T1- and T2-maps remove the strong SNR decay associated with the use of the EC. It is a fact that these strong intensity variations reduce the way the image is ridden compare to uniform signal intensity (provided by a QVBC for example). The T1 and T2 relaxation times obtained on the colon walls present an important standard deviation from the mean value. This is due to heterogeneity and size of the *in vivo* structures but also depends on the fit quality. Theoretically, in each pixel it is possible to obtain a value of the T1 and T2. Practically, some fits present a coefficient of determination too low to be taken into account. A part of the problem is due to the high and rapid variation of structures observed *in vivo*.

477 Moreover, structures are very thin and can be essentially made of fat (for deeper structures for
478 example). The non-significant values are removed from the results and can cause holes in the
479 maps. A further study will be to evaluate the influence of the number and values of flip angles
480 on fit quality. The 0.98 threshold here is chosen following what anatomical structures need to
481 be seen in the EC FOV. It corresponds to a setting neither too general nor drastic. Information
482 furnished by a fit quality of 0.98 are corresponding to the colon wall and surrounding structures.
483 When designing the coil, there were no considerations of heating and/or specific absorption
484 rate (SAR) taken into account. Indeed, during MR procedures, the radiofrequency power
485 transmitted (from the emission coil) for imaging is turned into heat through resistive losses.
486 Furthermore, when a conductor is placed within the imaging volume (as with the use of ECs)
487 the RF electrical field accompanying the RF magnetic field pulses induces currents that can
488 cause local concentration of the SAR [25, 26, 52, 53]. In our case, the loop can be considered
489 as a wire but the quarter wavelength (at 200 MHz) is much higher than the length of the loop.
490 It has been assumed that wire lengths less than a quarter wavelength are generally safe.
491 The presence of an active decoupling circuit is also mandatory to reduce SAR
492 concentration [24]. Further work will be to combine the EC with magnetic resonance
493 spectroscopy (MRS) in the colon wall. MRS could provide information on the biochemical
494 composition of deep tissue layers. Quantitative analysis of the biochemical contents could
495 also bring new markers to improve the measurements. But several challenges still lie ahead,
496 such as strong B_0 inhomogeneities and phase shift due to the difference of magnetic
497 susceptibility between air in the lumen and wall tissues.

498 In order to exploit the described work, a longitudinal follow up over six months involving
499 24 mice will be performed to explore *in vivo* the gastrointestinal wall in its entire depth. MR
500 ECs open new perspectives as a minimally invasive technique to characterize colorectal

501 cancer lesions and inflammatory bowel disease and then, offer a new technique to better
502 understand the development of the colon wall pathologies.
503

504 **CONCLUSION:**

505 In conclusion, an endoluminal coil was developed for imaging the colon and rectum walls on
506 a mouse model of colitis. The strong local SNR increase provided by the use of such coils
507 allows imaging the gastrointestinal tract at very high spatial resolution (up to approximately
508 40 μm in plane resolution with a slice thickness of 234 μm). Associated to a dedicated
509 protocol, it provides details in deep still thin structures of colon wall.

510

511 **Author's Contribution:**

512

513 DOREZ: Protocol/project development, data collection or management and data analysis

514 SABLON: Protocol/Project development, data collection

515 CANAPLE: Protocol/project development

516 SAINT-JALMES: Protocol/project development

517 GAILLARD: Data collection

518 MOUSSATA: Protocol/project development, data collection or management

519 BEUF: Protocol/project development, data collection or management

520

521 **COMPLIANCE WITH ETHICAL STANDARDS:**

522 **Disclosure of potential conflicts of interest:** the authors declare that they have no conflict of
523 interest.

524

525 **Research involving animals:** all applicable international, national, and/or institutional
526 guidelines for the care and use of animals were followed.

527 All procedures performed in studies involving animals were in accordance with the ethical
528 standards of the institution or practice at which the studies were conducted.

529

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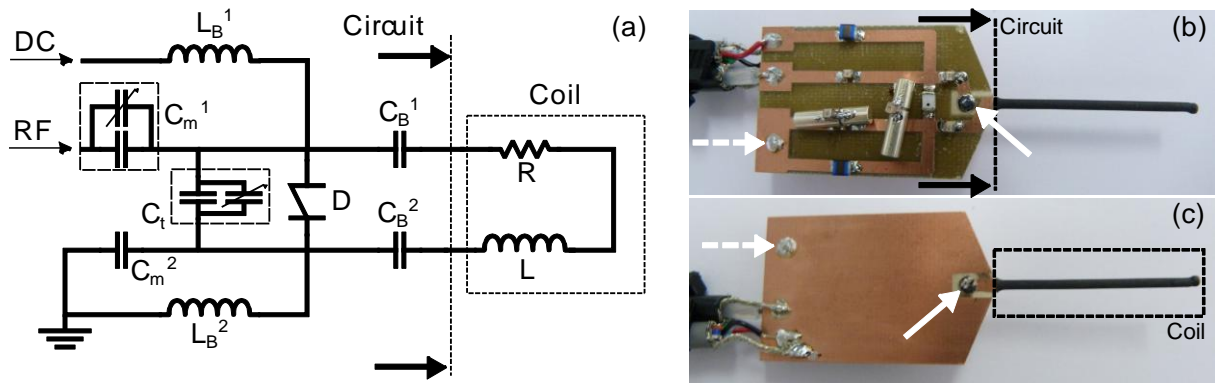
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658 LIST OF FIGURES

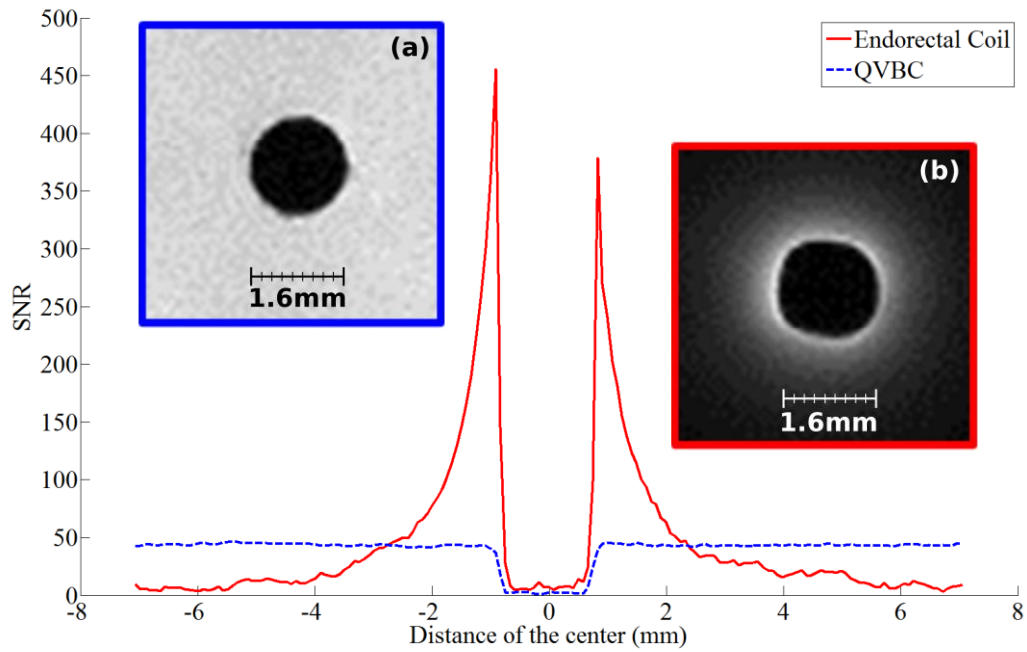


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660 Legend of Fig1:

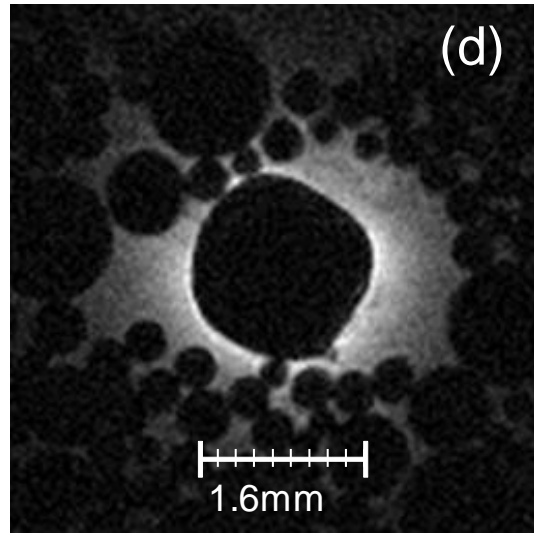
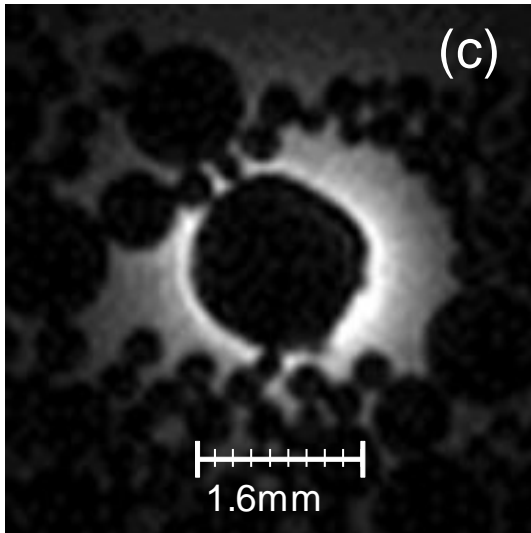
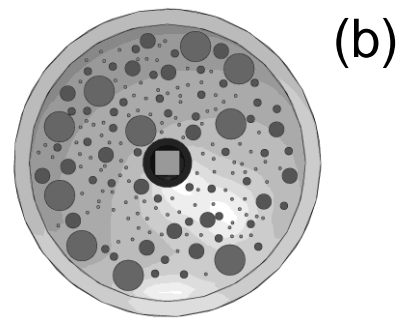
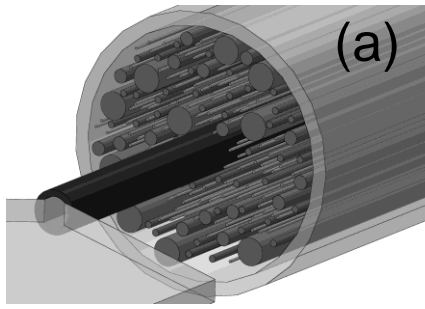
661 **Fig 1** (a) Electric scheme of the coil. The corresponding part between the scheme and the pictures are
 662 illustrated. (b) The upper face of the coil with the soldered components constitutes the RLC resonator.
 663 On (c) is represented the lower face of the prototype. Straight white arrow is the location of the plated
 664 drill between the two faces for connecting the electrical circuit to the coil. White dashed arrow is the
 665 location of the connection between the grounds (DC and RF) to the ground plane. Values of
 666 components are mentioned in Table 1

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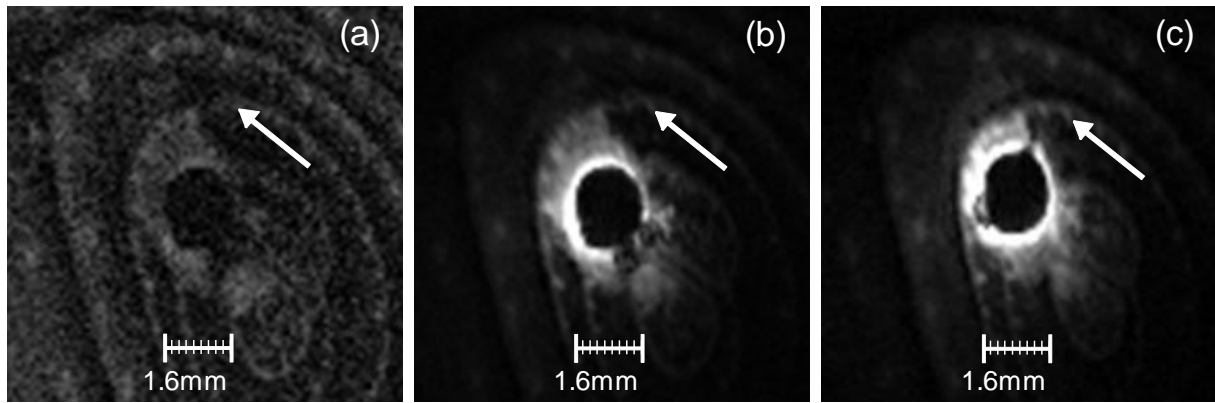
Legend of Fig 2:

Fig 2 Comparison of the EC SNR profile (straight profile) and the QVBC SNR profile (dashed profile) obtained on a phantom containing 1.25 g/L NiSO₄ and 5 g/L NaCl. (a) is the image used for plotting the QVBC profile and (b) the image used for the EC profile. Close to the coil the SNR is drastically better than the one obtained with a dedicated quadrature birdcage coil



Legend of Fig 3:

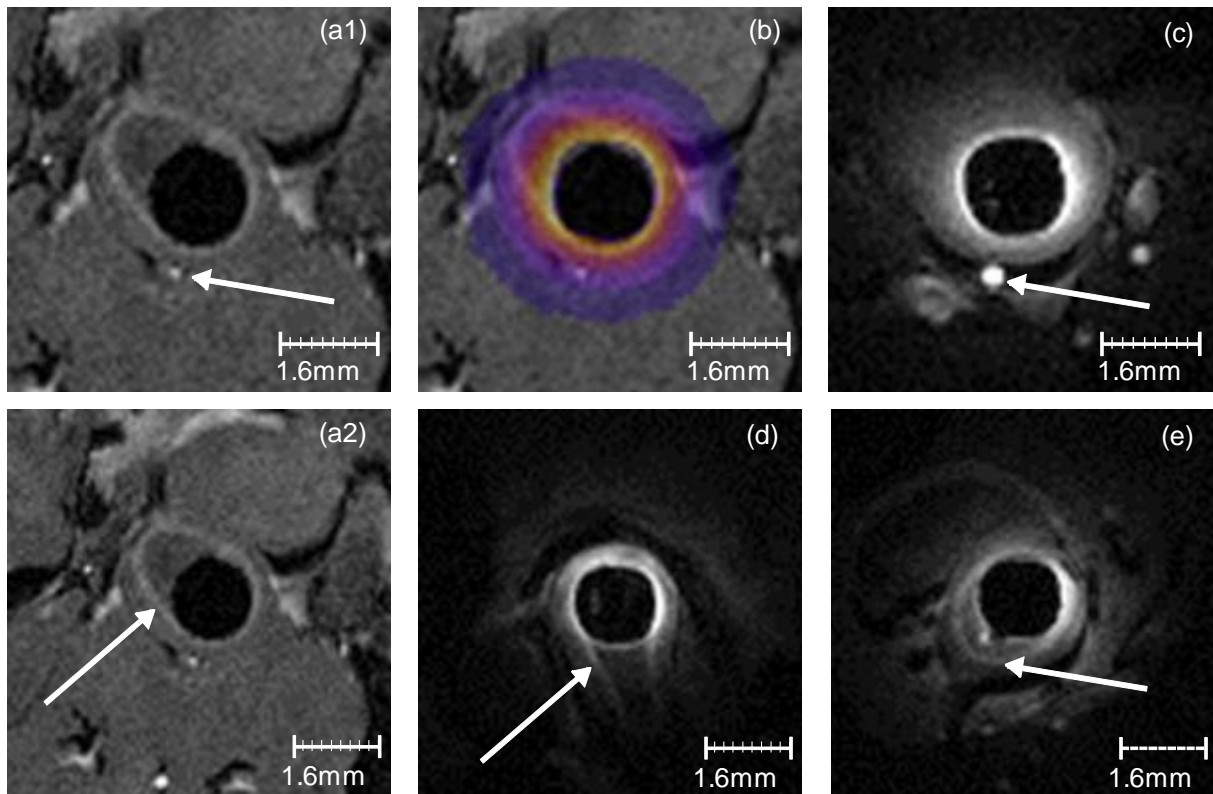
Fig 3 Images of the phantom created for the purpose of the experiments. (a) and (b) represent a reconstruction of the phantom containing optical fibers of different diameters. (c) and (d) are the corresponding MR images obtain with FLASH sequences at different voxel size ($50 \times 50 \mu\text{m}^2$ for (c) and $39 \times 39 \mu\text{m}^2$ (d)). The slice thickness was kept constant and equals to $234 \mu\text{m}$



Legend of Fig 4:

Fig 4 MR images of onion. (a) the volume coil was set in E/R mode, the lamellar structure represented by the white arrow are not correctly distinguishable. Switching the configuration by setting the VC in emission mode and the EC in reception mode the SNR and image quality are improved. On (b) the voxel size was decrease up to $78 \times 104 \mu\text{m}^2$ lamellar surface can be depicted more easily. Finally, the voxel size was reduced to $39 \times 39 \mu\text{m}^2$ on (c). Along the three MR images the slice thickness was set to $234 \mu\text{m}$

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Legend of Fig 5:

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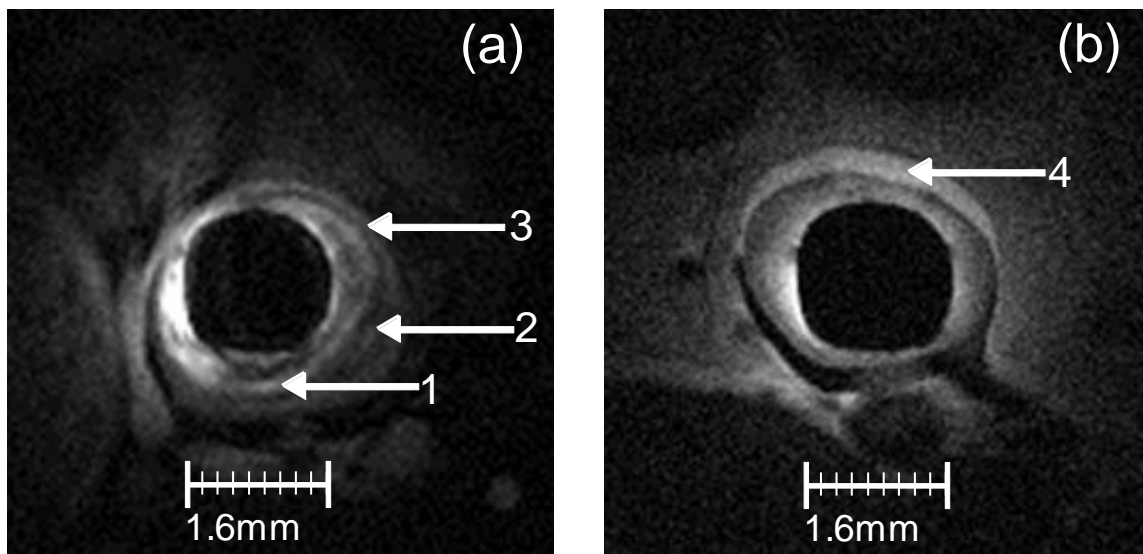
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Fig 5 *in vivo* experiments were performed on a mouse model of colitis. MR images acquired with the QVBC show blood vessels on (a1) and the colon wall on (a2) – white arrows. The SNR is too low to correctly distinguish the colon wall layers in details. The use of the EC allows to increase the local SNR. On (b) is represented the ratio between the SNR ratio of the QVBC and the EC. The EC FOV is sufficient for the region of interest. On (c), white arrow locates the same blood vessel than on (a1) and it is used for colocalization. (d) and (e) are showing the colon wall layer on different slices

701



702

703 Legend of Fig 6:

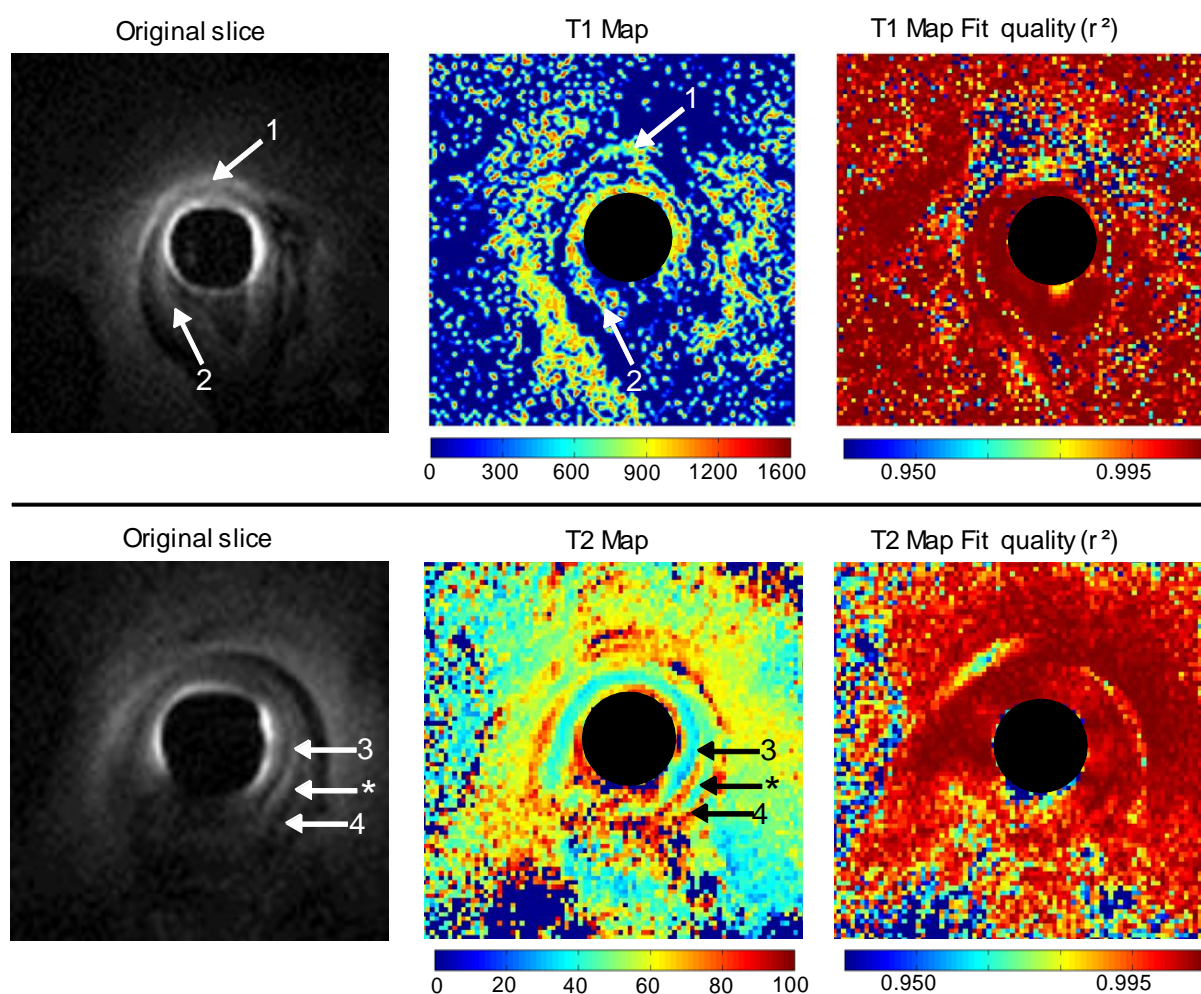
704 **Fig 6** From (a) to (b) the spatial resolution was increased up to $39 \times 39 \mu\text{m}^2$ (slice thickness of $234 \mu\text{m}$).

705 It is then possible to depict the colon wall complex in details. (1) refers to the mucosa, (2) to the

706 submucosa and (3) to the muscularis propria. (4) Represents a thin muscle layer closed to the coil. On

707 (b) the colon wall seems to have disappeared

708



710
711 Legend of Fig 7:

712 **Fig 7** T1- and T2-maps obtained on different slices. (1) indicates the location of a thin muscle
713 structure having T1 relaxation times lower than the ones observed on the colon wall (2). (3) and (4)
714 represents two different layers (mucosa and muscularis propria) which nearly identical physiological
715 constitutions so on with their respective T2 relaxation times. (*) locates the layer between the two
716 previous one and show a very different T2 time

717

718 TABLES

719

Table 1 Components used to build the endoluminal coil with values, manufacturers and references.

Function	Name	Value	Components
Tuning	Ct	22pF + 1pF	Non-variable non-magnetic capacitor
		from 0.8pF to 8pF	Non-magnetic trimmer capacitor
Matching	Cm1	5.6pF	Non-variable non-magnetic case A capacitor
		from 0.8pF to 8pF	Non-magnetic trimmer capacitor
	Cm2	5.6pF + 2.2pF + 1pF	Non-variable non-magnetic case A capacitor
Blocking	Cb1	100pF + 82pF	Non-variable non-magnetic case A capacitor
	Cb2	100pF + 82pF	Non-variable non-magnetic case A capacitor
Active decoupling	D	X	PIN diode
	LB1	1.2μH	Choke inductors
	LB2	1.2μH	Choke inductors

720

721

Table 2 Sequence and parameters used for the *in vivo* protocol. The scout sequences are not mentioned.

Sequence	Orientation	FOV (mm ²)	Slab/Slice Thickness (mm)	TR/TE (ms/ms)	Flip Angle (°)	Matrix	Voxel Size (μm ³)	Receive BW (kHz)	Acquisition Time
Flash 3D	Axial	16x16	1x15	20/6.653	15/20/25	192x192x64	83x83x234	25	8: 11'
FLASH 3D	Coronal	32x32	1x5	23.87/8.19	20	384x384x16	83x83x312	25	2: 26'
Multi TE	axial	10x10	6x1.5	2000/12,8+6x12,6	/	128x128x66	78x78x308	25	6: 24'

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723

Table 3 Sequences used for the characterization of the EC and the assessment of the EC during *in vitro* experiments.

Sequence	Orientation	FOV (mm ²)	Slab Thickness (mm)	TR/TE (ms/ms)	Flip Angle (°)	Matrix	Voxel Size (μm ³)	Receive BW (kHz)	Acquisition Time
Flash 3D	Axial	16x16	1x15	20/6.653	20	192x192x64	83x83x234	25	8: 11'
FLASH 3D	axial	10x10	1x15	20/6.1	20	256x256x64	39x39x234	25	10: 55'

724